DYCE 41.12. BCAD VI 61.721.5004 3:44:43 bW [E934611 D9A||6]µ4 [11.06]. 2AB: ABBL-110. DNIS: 81.73300 93. CSID: 205 281 2016 D14:09 |

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CLAIMS

What is claimed is:

1-33. (canceled)

34. (previously amended) A method for screening for genes whose expression is altered by disease, age, or exogenous agent, comprising:

screening a sample microarray comprising genes from a library, cells or animal exposed to the disease, age or exogenous agent, wherein expression of all of the genes is under control of the same regulatory element; and

comparing the expression of the genes to expression of control genes from a library, cells or animal not exposed to the disease, age or exogenous agent.

35. (currently amended) The method of claim 34 A method for screening for genes whose expression is altered by disease, age, or exogenous agent, comprising:

screening a sample microarray comprising genes from a library, cells or animal exposed to the disease, age or exogenous agent, wherein expression of all of the genes is under control of the same regulatory element; and

comparing the expression of the genes to expression of control genes from a library, cells or animal not exposed to the disease, age or exogenous agent;

wherein the microarray further comprises control genes that are not under the control of the same regulatory element.

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36. (currently amended) The method of claim 34 A method for screening for genes whose

expression is altered by disease, age, or exogenous agent, comprising:

screening a sample microarray comprising genes from a library, cells or animal

exposed to the disease, age or exogenous agent, wherein expression of all of the genes is under

control of the same regulatory element; and

comparing the expression of the genes to expression of control genes from a library.

cells or animal not exposed to the disease, age or exogenous agent;

wherein the regulatory element is selected from the group of regulatory elements

consisting of osmotic response element, retinoic acid response element, conserved proximal

sequence element, vitamin D response element, sterol response element, TNF-alpha response

element, serum response element, cAMP response element, antioxidant response element,

glucotocorticoid modulatory element, gonadotropin-releasing hormone-response element,

pheromone response element, insulin response element, interferon consensus response element.

estrogen response element, hypoxia response element, E2F transcription factor, xenobiotic response

element, endoplasmic reticulum stress response element, iron-response element, androgen response

element, stress response element, RAS-responsive element binding protein 1, and transforming

growth factor, beta-1 response element.

37. (canceled)

38. (original) The method of claim 34 wherein the disease is selected from the group

consisting of neurological disorders, cardiovascular disorders, bone and muscle disorders, blood or

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circulation related disorders, and cancer.

39. (original) The method of claim 38 wherein the diseases are selected from the group

consisting of Alzheimer's disease, Parkinson's disease, Huntington's disease, myocardial

hypertrophy, atherosclerosis, myocardial infarction, osteoarthritis, osteoporosis, and autoimmune

disorders.

40. (original) The method of claim 38 wherein the cancers are selected from the group

consisting of breast cancer, prostatic hypertrophy, prostatic cancer, colon cancer, chronic

lymphocytic leukemia, acute lymphocytic leukemia, brain tumors, pancreatic cancer, and

heptatomas.

41. (canceled)

42. (previously added) The method of claim 34 wherein the exogenous agent is a drug or

toxin.

43. (previously added) The method of claim 34 wherein the library is derived from cells

or tissues treated with one or more compounds in vitro.

44. (previously added) The method of claim 34 wherein the library is derived from cells

obtained from an individual of a particular age, having a particular disease or disorder, or derived

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from the neurological system, the cardiovascular system, the musculoskeletal system, or cancerous

tissues.

45. (previously added) The method of claim 34 wherein the exogenous agent is selected

from the group consisting of proteins or peptides, sugars or polysaccharides, nucleic acid molecules,

and synthetic molecules.

46. (previously added) The method of claim 43 wherein the compound is selected from

the group consisting of proteins or peptides, sugars or polysaccharides, nucleic acid molecules, and

synthetic molecules.

47. (new) The method of claim 35 wherein the disease is selected from the group

consisting of neurological disorders, cardiovascular disorders, bone and muscle disorders, blood or

circulation related disorders, and cancer.

48. (new) The method of claim 47 wherein the diseases are selected from the group

consisting of Alzheimer's disease, Parkinson's disease, Huntington's disease, myocardial

hypertrophy, atherosclerosis, myocardial infarction, osteoarthritis, osteoporosis, and autoimmune

disorders.

49. (new) The method of claim 47 wherein the cancers are selected from the group

consisting of breast cancer, prostatic hypertrophy, prostatic cancer, colon cancer, chronic

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lymphocytic leukemia, acute lymphocytic leukemia, brain tumors, pancreatic cancer, and

heptatomas.

50. (new) The method of claim 35 wherein the exogenous agent is a drug or toxin.

51. (new) The method of claim 35 wherein the library is derived from cells or tissues

treated with one or more compounds in vitro.

52. (new) The method of claim 35 wherein the library is derived from cells obtained from

an individual of a particular age, having a particular disease or disorder, or derived from the

neurological system, the cardiovascular system, the musculoskeletal system, or cancerous tissues.

53. (new) The method of claim 35 wherein the exogenous agent is selected from the

group consisting of proteins or peptides, sugars or polysaccharides, nucleic acid molecules, and

synthetic molecules.

54. (new) The method of claim 35 wherein the compound is selected from the group

consisting of proteins or peptides, sugars or polysaccharides, nucleic acid molecules, and synthetic

molecules.

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